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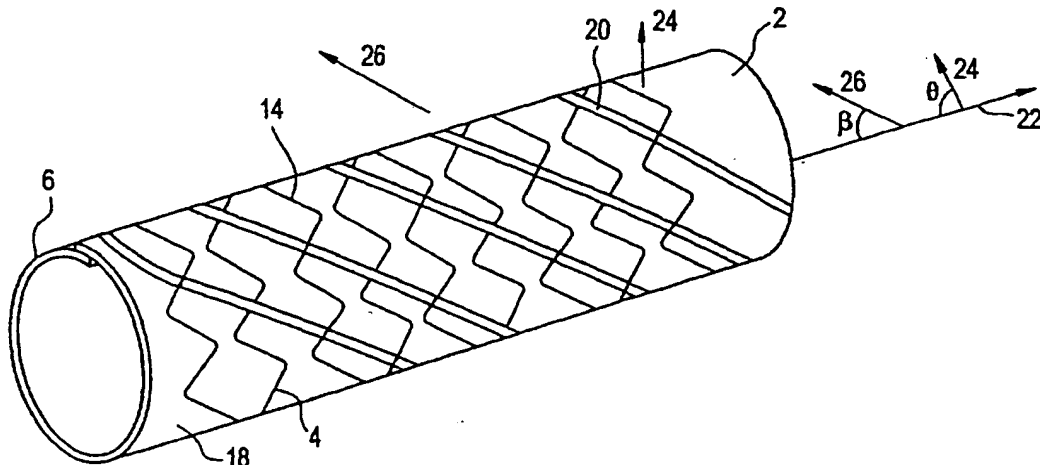
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(54) Title: STENT-GRAFT WITH HELICALLY ARRANGED SECUREMENT MEMBER



(57) Abstract: An improved endoluminal prosthesis is described which possesses increased flexibility and longitudinal compliance with a stent/graft/stent securement member combination.

STENT-GRAFT WITH HELICALLY ARRANGED SECUREMENT MEMBER

FIELD OF THE INVENTION

The present invention relates to an implantable endoluminal prosthesis. More particularly, the present invention relates to a stent-graft endoluminal prosthesis offering increased flexibility and compliance.

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BACKGROUND OF THE INVENTION

An endoluminal prosthesis is a medical device commonly known to be used in the treatment of diseased blood vessels. An endoluminal prosthesis is typically used to repair, replace, or otherwise correct a damaged blood vessel. An artery or vein may be
10 diseased in a variety of ways. An endoluminal prosthesis is therefore designed to be used to prevent or treat a wide variety of defects such as stenosis of the vessel, thrombosis, occlusion, or an aneurysm.

One type of endoluminal prosthesis used in the repair of diseases in various body
15 vessels is a stent. A stent is a generally longitudinal tubular device formed of biocompatible material which is useful to open and support various lumens in the body. For example, stents may be used in the vascular system, urogenital tract and bile duct, as well as in a variety of other applications in the body. Endovascular stents have become widely used for the treatment of stenosis, strictures, and aneurysms in various blood
20 vessels. These devices are implanted within the vessel to open and/or reinforce collapsing or partially occluded sections of the vessel.

Stents are generally open ended and are radially expandable between a generally unexpanded insertion diameter and an expanded implantation diameter which is greater
25 than the unexpanded insertion diameter. Stents are often flexible in configuration, which allows them to be inserted through and conform to tortuous pathways in the blood vessel. The stent is generally inserted in a radially compressed state and expanded either through a self-expanding mechanism, or through the use of balloon catheters.

A graft is another type of endoluminal prosthesis which is used to repair and replace body vessels. Whereas a stent provides structural support to hold a damaged vessel open, a graft provides an artificial lumen through which blood may flow. Grafts are tubular devices which may be formed of a variety of materials, including textile and non-textile materials. One type of non-textile material particularly suitable for use as an implantable prosthesis is polytetrafluoroethylene (PTFE). PTFE exhibits superior biocompatibility and low thrombogenicity, which makes it particularly useful as vascular graft material in the repair or replacement of blood vessels. In vascular applications, the grafts are manufactured from expanded PTFE (ePTFE) tubes. These tubes have a microporous structure which allows natural tissue ingrowth and cell endothelialization once implanted in the vascular system. This contributes to long term healing and patency of the graft.

It is also known to combine a stent and a graft to form a composite medical device. Such a composite medical device provides additional support for blood flow through weakened sections of a blood vessel. In endovascular applications the use of a stent/graft combination is becoming increasingly important because the combination not only effectively allows the passage of blood therethrough, but also ensures the implant will remain open.

Several types of stent/graft inventions are known in the art. U.S. Patent No. 5,151,105 issued to Kwan-Gett discloses a collapsible textile vessel sleeve with stent members positioned at opposite ends of the sleeve. The device is specifically designed to provide a vessel sleeve that is collapsible to a very small diameter in order that it may be placed in position within the abdominal or thoracic aorta by a catheter via the lumen of the femoral artery. Such a procedure obviates the need for a major surgical intervention, and reduces the risks associated with such a procedure.

Other stent/graft composite devices using a textile fabric are shown in U.S. Patent No. 5,628,788 to Pinchuck.

As mentioned above, ePTFE may also be used as graft material in stent/graft endoprostheses. One example of an ePTFE stent/graft device is shown in U.S. Patent No. 5,700,285 issued to Myers, et al. Myers discloses a tubular intraluminal graft in the form of a tubular diametrically adjustable stent having an interior and exterior tubular covering of porous expanded polytetrafluoroethylene. The tubular covering surrounds the stent so that the stent is contained during contraction and expansion in the delivery process.

Stents are effectively used in combination with grafts as the composite endoluminal prosthesis allows blood flow through the vessel created by the graft, while the stent maintains its patency. It has been found however that stent/graft composite devices exhibit reduced flexibility and longitudinal compliance. Longitudinal compliance is of particular importance to such stent/graft endoluminal prosthesis as the device must be intraluminally delivered through tortuous pathways of a blood vessel to the implantation site where the stent is expanded. The increased width creates a profile of increased size of the prosthesis which may present problems in placing and expanding the prosthesis in smaller arteries and veins which demand a smaller, more flexible endoluminal prosthesis. Such reduction of compliance and flexibility is caused by the increased thickness of the composite device and also by the technique used to secure the stent to the graft.

In order to solve such problems, several stent/graft devices have been developed. International publication number WO 97/21403 to Prograft discloses a stent graft combination comprising a stent member with an inner and outer surface, a tubular graft member, and a ribbon covering only a portion of at least one of the inner and outer surfaces of said stent member for securing the stent member and graft member to one another. This device uses a broad ribbon in order to increase the potential bonding surface area between the coupling member and the graft member to enhance the structural integrity of the stent/graft device while reducing the total thickness of the composite graft. The coupling ribbon used in the Prograft stent/graft device essentially tracks the path of the stent in coupling the stent to the graft, i.e., the coupling ribbon and

the stent are helically wound to form a spiral configuration where both have the same angular orientation with respect to the longitudinal axis of the graft.

5 The use of such a broad coupling member, however, may present a variety of problems. The use of such a broad coupling member also may decrease the overall flexibility of the prosthesis. The use of a broad coupling member may furthermore, increase radial stiffness of the prosthesis. In addition wrapping the ribbon about the stent at the same angular orientation as the stent tends to decrease the flexibility and expandability of the stent.

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It is desirable to provide a stent/graft composite device which secures the stent to the graft without exhibiting a significant reduction in longitudinal compliance or flexibility.

15 SUMMARY OF THE INVENTION

It is an advantage of the present invention to provide an endoluminal prosthesis including a stent/graft composite device having increased flexibility.

20 It is also an advantage of the present invention to provide a stent/graft composite device where the stent is secured to the graft so as to not reduce the flexibility and longitudinal compliance of the device.

25 The present invention provides a thin suture, or filament-like securement member which secures the stent to the graft by being helically arranged at an angle with respect to the longitudinal axis of the graft, such that the angle does not equal the angle formed with respect to the helical windings of the stent. The present invention is designed to be able to provide structural integrity for the prosthesis while also substantially reducing its size. The prosthesis exhibits a reduced profile and increased flexibility over prior art. Furthermore, by angularly orienting the wrap with respect to the stent, the radial wrap
30 provides the prosthesis with a more efficient means of securing the stent cohesively together. The angular orientation in regard to the stent, further provides a more efficient means of securing the stent to the tubular member.

In attaining these and other advantages, the present invention provides an endoluminal prosthesis comprising a polytetrafluoroethylene tubular member with a luminal surface, an exterior surface, and a longitudinal axis. A diametrically deformable stent is circumferentially disposed on the luminal or exterior surface of said tubular member, said stent being formed from an elongate wire helically wound defining a plurality of spaced apart windings disposed at a first angle with respect to said longitudinal axis, and an elongate securement member securing said stent to said tubular member, said securement member being helically arranged at a second angle with respect to said longitudinal axis so that said first angle is not congruent to said second angle.

A method of making the endoluminal prosthesis of the present invention is also disclosed. The method comprises providing a polytetrafluoroethylene tubular member, said tubular member comprising a luminal surface, an exterior surface, and a longitudinal axis. The method of making the prosthesis further comprises disposing a diametrically deformable stent circumferentially on the luminal or exterior surface of said tubular member, said stent being formed from an elongate wire helically wound defining a plurality of spaced apart windings disposed at a first angle with respect to said longitudinal axis; and helically wrapping an elongate securement member to secure said stent to said tubular member, said elongate securement member being wrapped at a second angle with respect to said longitudinal axis, so that said second angle is not congruent to said first angle of said stent.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a perspective showing a collapsed wave-like stent which may be used in the present intraluminal prosthesis.

Figure 2 is a perspective showing a stent similar to the stent of Figure 1 expanded about an exterior surface of an ePTFE graft tube.

Figure 3 is a perspective showing a composite endoluminal prosthesis comprising another stent embodiment disposed on the exterior surface of a tubular member.

Figure 4 is a perspective showing of the endoluminal prosthesis of Figure 3 including a securement member for supporting the stent to the graft.

Figure 5 is a perspective showing a further embodiment of the present invention with the securement member angularly oriented at two different angles to secure the stent to the tubular member.

Figure 6 is a perspective showing of a still further embodiment of an endoluminal prosthesis of the present invention wherein the securement member forms loops around elbows of the stent.

Figure 7 is a perspective showing of yet another embodiment of an endoluminal prosthesis of the present invention wherein the securement member loops around leg segments of the stent.

Figure 8 is a perspective showing another embodiment of the endoluminal prosthesis of the present invention wherein a nested stent is shown with a securement member securing said stent at its elbows.

DETAILED DESCRIPTION OF THE INVENTION

The following is a detailed description of the preferred embodiments of the present invention. The description is meant to describe preferred embodiments, and is not meant to limit the invention in any way.

The present invention provides an improved endoluminal prosthesis comprising a polytetrafluoroethylene tubular member with a luminal surface, an exterior surface, and a longitudinal axis. A diametrically deformable stent is circumferentially disposed on the surface of the tubular member. The stent is formed from an elongate wire helically wound defining a plurality of spaced apart windings disposed at a first angle with respect

to said longitudinal axis. An elongate securement member secures the stent to said tubular member, the securement member is helically arranged at a second angle with respect to said longitudinal axis so that said first angle is not congruent to said second angle.

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The design allows securement of a stent to a graft in such a manner to ensure optimal properties of flexibility and reduced profile. Specifically, the helical angularly oriented direction of the securement member, which is wrapped circumferentially around the tubular member creates an angle with respect to the longitudinal axis of the tubular member which is incongruous to the angle at which the windings of the stent are disposed with respect to the longitudinal axis of the prosthesis. This allows for structural integrity of the prosthesis with a minimal amount of material comprising the securement member.

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Various stent types and stent constructions may be employed in the present invention. Among the various stents useful include, without limitation, self-expanding stents and balloon expandable stents. The stents may be capable of radially contracting, as well, and in this sense can best be described as radially distensible or deformable. Self-expanding stents include those that have a spring-like action which causes the stent to radially expand, or stents which expand due to the memory properties of the stent material for a particular configuration at a certain temperature. Nitinol is one material which has the ability to perform well while both in spring-like mode, as well as in a memory mode based on temperature. Other materials are of course contemplated, such as stainless steel, platinum, gold, titanium, and other biocompatible metals, as well as polymeric stents.

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Referring now to Figures 1 and 2 of the drawings, the composite prosthesis 5 of the present invention includes a stent 2 shown in Figure 1, as well as a similar stent 3 shown in Figure 2 of the drawings. Stent 2 is shown in a collapsed configuration in Figure 1. Stents 2 and 3 include a plurality of spaced apart windings 4 which include successive upper wave-like peaks 6 and lower wave-like peaks 8. The upper and lower wave-like peaks are connected via leg segments 10 of the stent. As seen in Figure 2,

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stent 3 shows parallel windings 4 arranged along the longitudinal axis of the prosthesis in such a manner that upper wave-like peaks 6 are nested within lower wave-like peaks 8 of adjacent windings along the stent. As used in this disclosure, the term nested refers to the stent configuration where successive upper wave-like peaks of the stent are
5 linearly aligned to fit within the successive lower wave-like peaks of the stent so that each wave is stacked (or nested) within the next adjacent winding. The stent shown in Figure 1 is more fully described in commonly assigned U.S. Patent No. 5,906,639 to Rudnick, et al., herein incorporated by reference.

10 Referring to Figure 2 of the drawings, the tubular graft member 12 of the endoluminal prosthesis is preferably formed of polytetrafluoroethylene (PTFE). In vascular applications, prostheses are most often manufactured from expanded PTFE (ePTFE) tubes. As a result of the stretching and expansion of the PTFE material, these tubes have a microporous structure which allows natural tissue ingrowth and cell
15 endothelialization once implanted in the vascular system. This contributes to long term healing and patency of the prosthesis. Tubular member 12 may be extruded as a tube or may be formed from an extruded sheet which is subsequently formed into a tubular structure. Textile or fabric constructions formed of PTFE or ePTFE yarns, filaments, or mesh may also be employed.

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As shown in Figure 2, stent 3 is applied in a radially expanded condition about the exterior surface of tubular graft member 12 to form a stent/graft composite of the type well known in the art. While the preferred embodiments described herein show stent 2 supported about the exterior surface of tubular graft member 12, it is within the
25 contemplation of the present invention to support the stent on the opposed luminal surface thereof.

Referring now to Figures 3 and 4, a composite stent/graft device 17 is shown. Device 17 includes a stent 14 positioned about an ePTFE tube 18. In the present
30 example tube 18 is formed from a sheet of ePTFE. Stent 14 is similarly formed of spaced apart windings comprising successive upper wave-like peaks and lower wave-like peaks connected via leg segments of the stent. As can be seen in Figure 3, the

upper wave-like peaks meet lower wave-like peaks of successive windings to form elbows of coincidence 16.

As mentioned above, diametrically deformable stent 14 is helically wound to
5 define a plurality of spaced apart windings 15. Windings 15 are disposed at a first angle with respect to the longitudinal axis of the ePTFE tubular member. The longitudinal axis of tubular member 18 is shown with directional arrow 22. Successive windings 15 of stent 14 are positioned at a first angle (as shown with directional arrow 24) θ with respect to the longitudinal axis of tubular member 18.

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The present invention provides a securement member 20 which is most preferably a flat, thin suture. Tape, thread, ribbon, or other elongate members may also be used. The securement member may be formed of a variety of materials. Securement member 20 is preferably less than 0.60 mm wide. The securement member is preferably
15 made from a textile material. In using the term textile material in this disclosure, it is meant to indicate any material which may be used to combine with other pieces of the same material to become part of a larger piece of fabric. Some materials which may be used as the securement member include, but are not limited to PTFE, ePTFE, Polyethylene terephthalate (PET), Polyether ether ketone (PEEK), polypropylene, fluorinated ethylene propylene (FEP), nylon (polyamide), polyurethane (PU) and
20 polyimide (PI), polybutylene terephthalate (PBT), polyurethane rubber (PUR), silicone and silicone rubber, and bio-absorbable materials, including poly(glycolic acid) (PGA), poly(lactic acid) (PLA).

25 As shown in Figure 4 of the drawings, securement member 20 is helically arranged at a second angle β which is non-congruent to said first angle θ . This design construction allows the prosthesis to be thin-walled, compliant, and more flexible, because it provides structural integrity using less covering in the form of securement member 20. When securement member 20 is angled at a different angle (as compared to
30 angle θ between the stent windings and axis) to longitudinal axis 22, it allows the securement of the stent to the graft using less material than previously used in prostheses of this type. More specifically, the angular arrangement of the securement member

allows a thin securement member, which allows for a more flexible, and thinner composite prosthesis. Both angle θ and angle β may equal any value from 0° to 180° with respect to longitudinal axis 22.

As mentioned above, a stent is preferably mounted on the exterior surface of tubular member 18. Securement member 20 is then helically wrapped around tubular member 18 at angle β with respect to longitudinal axis 22 of tubular member 18. Securement member 20 may be wrapped completely exteriorly of stent 4, or it may be interwoven between stent 4 and the exterior surface of tubular member 18. Securement member 20 is then adhered to tubular member 18. Adherence is typically accomplished by sintering the composite prosthesis. Sintering, as used in the present disclosure means heating the composite prosthesis to a temperature below its melting point, yet sufficient to thermally adhere the prosthesis. The heat of sintering differs for different materials. An adhesive may be used with the sintering process or the securement member may be adhered with an adhesive without sintering.

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The securement member 20 ensures the integrity of the composite prosthesis. The securement member is typically adhered to the tubular member in some manner, and may weave in and out of the leg segments of the stent in order to securely attach the stent. The term adhered as used in this disclosure refers to the attaching of the securement member to the tubular member in any manner. It includes without limitation, lamination, thermally adhering, sintering, RF welding, attaching with an adhesive, and any combination of the above. The securement member may be adhered noncontinuously at selected areas, or may be continuously adhered throughout its entire length.

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Referring now to Figure 5 of the drawings, securement member 20 may be wound at more than one angle with respect to the longitudinal axis 22. Directional arrows 26 and 28 show the different orientation (as well as the different angles they make as compared to the longitudinal axis 22) of the securement member with respect to longitudinal axis 22. When the securement member is orientated in two different directions with respect to the longitudinal axis as shown in Figure 5, the securement

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member may intersect other helically oriented strips of the securement member to form nodes of intersection 30. Securement member 20 may be adhered to itself at nodes 30, or may be adhered to itself, as well as the tubular member. Preferably securement member 20 is sintered to itself at the segment it intersects, as well as to the tubular prosthesis at node 30.

Figure 6 shows another embodiment of the prosthesis of the present invention. Similar to the stent in Figure 3, elbows of coincidence 16 are formed where the successive upper and lower wave-like peaks meet in successive stent windings. Thin suture 32 forms a loop around the peaks at the elbows of coincidence. In this manner, suture 32 forms eyelets 34 holding the successive peaks together at elbows 16. Suture 32 may be adhered to the tubular member at eyelets 34, preferably by lamination.

Figure 7 shows another embodiment of the present invention. Securement member consists of thin suture 32, which attaches stent 38 to tubular member 40 by forming loops 36 around leg segments 42 of stent 38. Suture 32 may be adhered to the tubular member along its length. The loops 36 are preferably adhered to the tubular member under leg segments 42 of the stent around which the loop is formed. The loops may also take the form of knots of different configurations and may be tied around the stent in many alternative methods.

Figure 8 shows yet another embodiment of the present invention. Stent 44 is in a nested configuration. Thin suture 32 forms loops 36 at wave-like peaks 46 in order to secure the stent to the tubular member. Similar to the prosthesis shown in Figure 7, the loops 36 are preferably adhered to the tubular member under the stent at loop 36.

Various changes to the foregoing described and shown structures would now be evident to those skilled in the art. Accordingly, the particularly disclosed scope of the invention is set forth in the following claims.

WHAT IS CLAIMED IS:

1. An endoluminal prosthesis comprising:
an elongate polytetrafluoroethylene tubular member having a luminal surface, an
5 exterior surface and extending along a longitudinal axis;
a diametrically deformable stent circumferentially disposed on the luminal or
exterior surface of said tubular member, said stent being formed from an elongate wire
helically wound defining a plurality of spaced apart windings disposed at a first angle
with respect to said longitudinal axis, and
10 an elongate securement member securing said stent to said tubular member, said
securement member being helically arranged at a second angle with respect to said
longitudinal axis, so that said first angle is non-congruent to said second angle.
2. The endoluminal prosthesis according to claim 1 wherein said securement
15 member is a substantially flat, thin suture.
3. The endoluminal prosthesis according to claim 2 wherein said securement
member is formed of a textile material, said material being selected from the group
consisting of PET, PTFE, and ePTFE.
20
4. The endoluminal prosthesis according to claim 1 wherein said securement
member secures said stent to said exterior surface of said tubular member.
5. The endoluminal prosthesis according to claim 1 wherein said securement
25 member is adhered to said tubular member.
6. The endoluminal prosthesis according to claim 5 wherein said securement
member is noncontinuously adhered to said tubular member at certain points.
- 30 7. The endoluminal prosthesis according to claim 5 wherein said securement
member is continuously adhered to said tubular member wherever said securement
member interfaces with said tubular member.

8. The endoluminal prosthesis according to claim 1 wherein said spaced apart windings of said stent include successive upper and lower wave-like peaks, and wherein said successive peaks are connected via leg segments of the stent.

5 9. The endoluminal prosthesis according to claim 8 wherein said successive upper and lower wave-like peaks of longitudinally adjacent windings of said stent are nested along the length of said longitudinal axis.

10 10. The endoluminal prosthesis according to claim 8 wherein said upper wave-like peaks of said windings meet said lower wave-like peaks of longitudinally adjacent windings to form elbows of coincidence between said adjacent windings.

15 11. The endoluminal prosthesis according to claim 10 wherein said securement member secures said stent at said elbows of coincidence by forming an eyelet securing said adjacent upper and lower peaks together.

12. The endoluminal prosthesis according to claim 8 wherein said securement member weaves in and out of said leg segments of said stent.

20 13. The endoluminal prosthesis according to claim 8 wherein said securement member secures said stent to said tubular member by forming loops around leg segments of said stent.

25 14. The endoluminal prosthesis according to claim 1 wherein said securement member is helically wound at said second angle with respect to said longitudinal axis, and further helically wound at a third angle with respect to said longitudinal axis, in such a manner that said third angle does not equal said second angle and said first angle.

30 15. The endoluminal prosthesis according to claim 14 wherein said second and third angles of said securement member are angularly oriented with respect to said longitudinal axis in such a manner that said securement member intersects itself at nodes of intersection as a result of said angular orientation.

16. The endoluminal prosthesis according to claim 15 wherein said securement member is adhered together and to the polytetrafluoroethylene tubular member at said nodes of intersection.

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17. A method of making an endoluminal prosthesis comprising:
providing a polytetrafluoroethylene tubular member, said tubular member comprising a luminal surface, an exterior surface, and a longitudinal axis;

10 disposing a diametrically deformable stent circumferentially on the luminal or exterior surface of said tubular member, said stent being formed from an elongate wire helically wound defining a plurality of spaced apart windings disposed at first angle with respect to said longitudinal axis; and

helically wrapping an elongate securement member to secure said stent to said tubular member, said elongate securement member being wrapped at a second angle
15 with respect to said longitudinal axis so that said second angle being non-congruent to said first angle of said stent.

18. The method according to claim 17 wherein said securement member is a substantially flat, thin suture.

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19. The method according to claim 17 wherein said securement member is formed of a spun web type of textile material, said material being selected from the group consisting of PET, PTFE, and ePTFE.

20. The method according to claim 17 wherein said securement member secures
25 said stent to said exterior surface of said tubular member.

21. The method according to claim 17 wherein said securement member is adhered to said tubular member.

30

22. The method according to claim 21 wherein said securement member is adhered to said tubular member noncontinuously at certain points.

23. The method according to claim 21 wherein said securement member is continuously adhered to the tubular member wherever said securement member interfaces with said tubular member.

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24. The method according to claim 17 wherein said securement member is adhesively adhered to said tubular member.

25. The method according to claim 24 wherein said securement member is adhesively adhered to said tubular member noncontinuously at certain points.

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26. The method according to claim 24 wherein said securement member is continuously adhesively adhered to said tubular member wherever said securement member interfaces with said tubular member.

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27. The method according to claim 17 wherein said stent is formed from an elongate wire helically wound with a plurality of longitudinally spaced turns including successive upper and lower wave-like peaks, and wherein said successive peaks are connected via leg segments of the stent.

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28. The method according to claim 27 wherein said securement member weaves in and out of said leg segments of said stent.

29. The method according to claim 17 wherein said securement member is helically wound at said second angle with respect to said longitudinal axis of said tubular member, and further helically wound at a third angle with respect to said longitudinal axis of said tubular member, said second and third angles being not equivalent.

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30. The method according to claim 29 wherein said second and third angles of said securement member are angularly oriented with respect to said longitudinal axis in such a manner that said securement member intersects itself at nodes of intersection as a result of said angular orientation.

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31. The method according to claim 30 wherein said securement member is adhesively adhered together at said nodes of intersection.

- 5 32. The method according to claim 31 wherein said securement member is adhesively adhered to the expanded polytetrafluoroethylene tubular member at said nodes of intersection.

FIG. 1

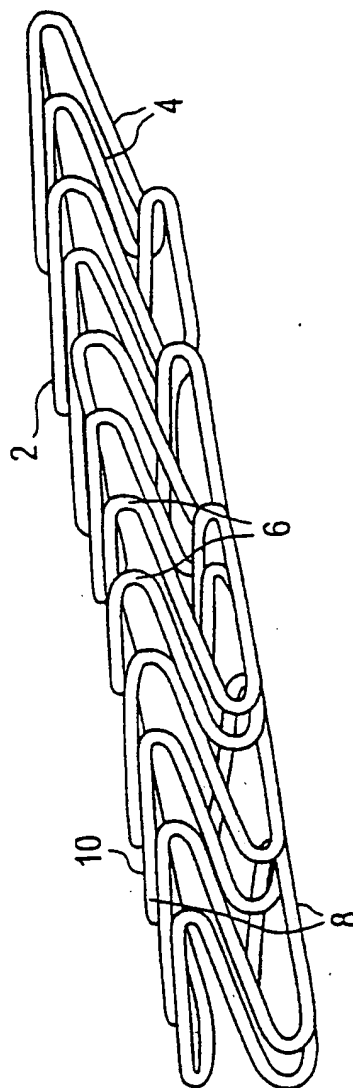


FIG. 2

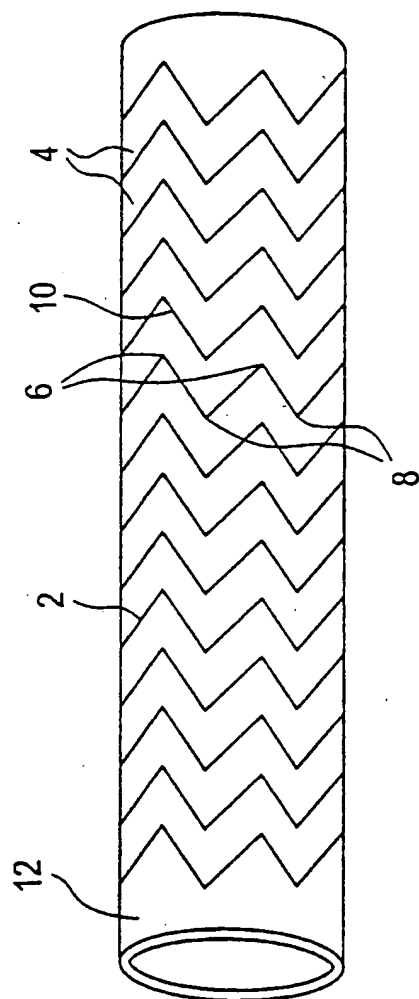


FIG. 3

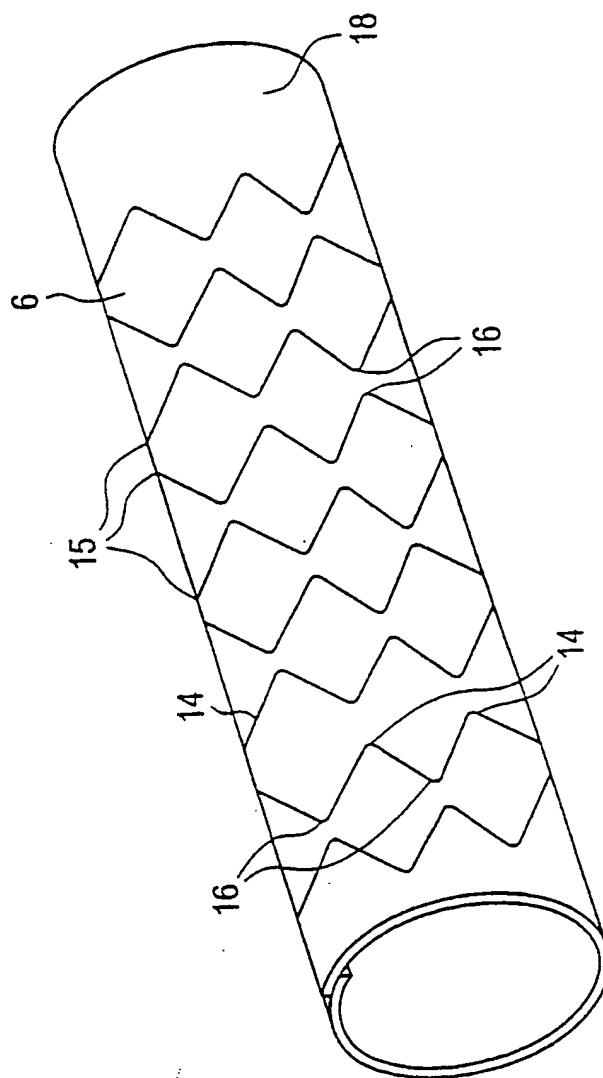
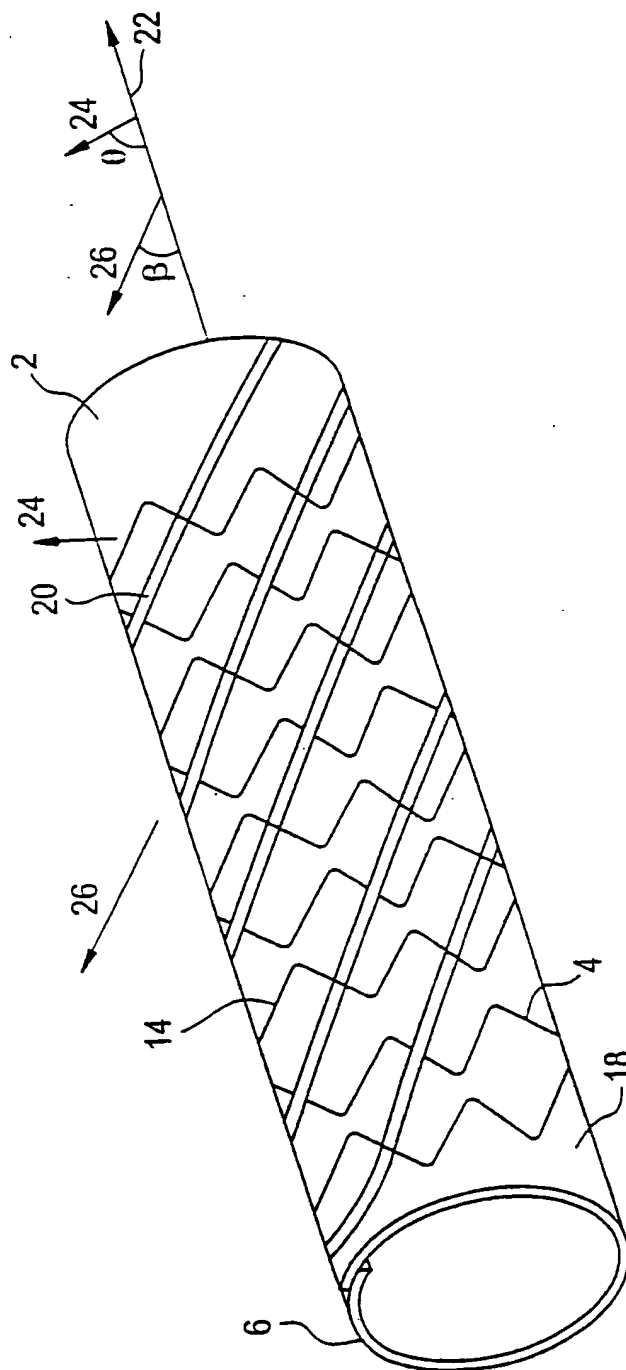


FIG. 4



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FIG. 5

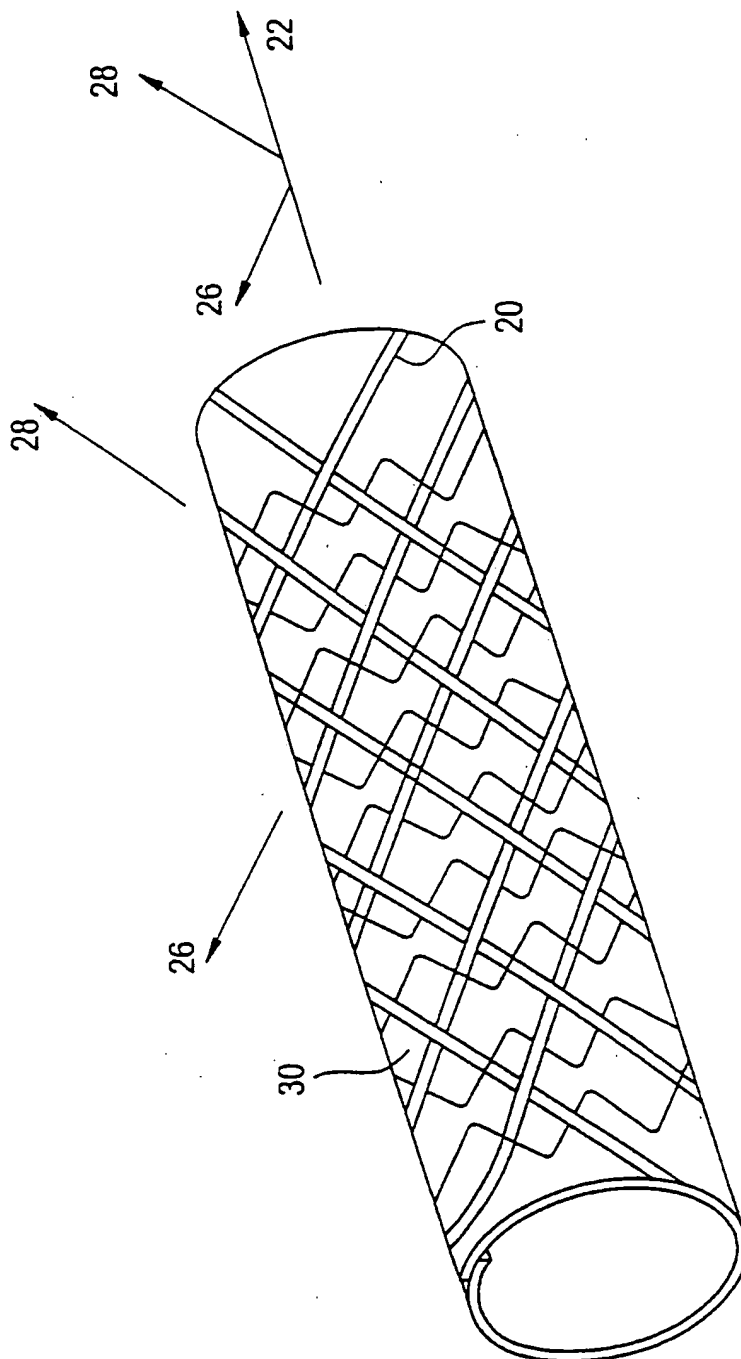


FIG. 6

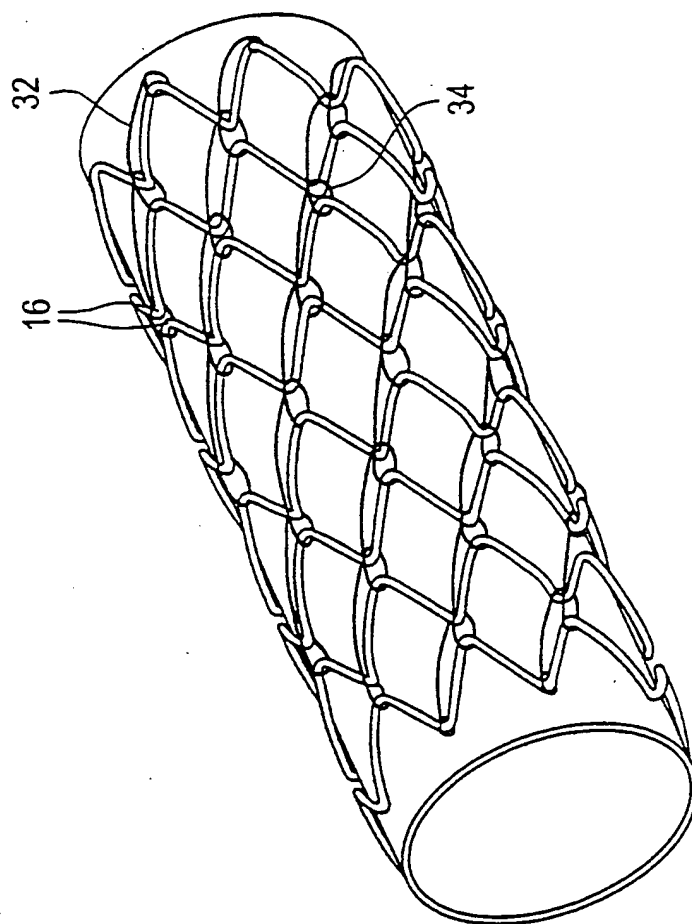


FIG. 7

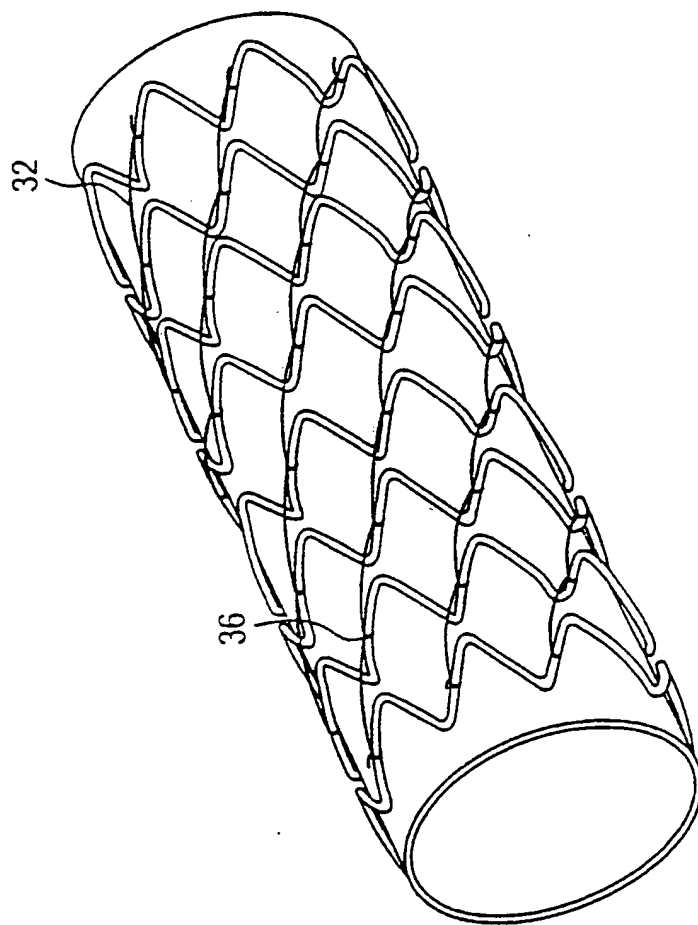
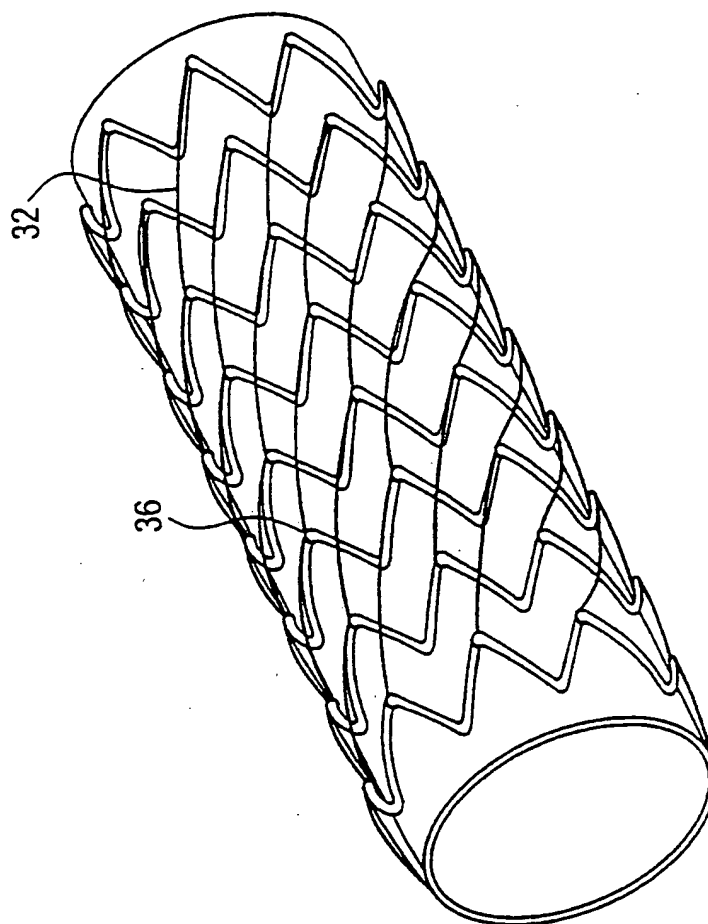


FIG. 8



INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 01/01261

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 A61F2/06

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 A61F

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	WO 98 27894 A (PROGRAFT MEDICAL INC) 2 July 1998 (1998-07-02) page 17, line 2 -page 19, line 24; figures 1-6	1,17
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